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**IN THE UNITED STATES PATENT AND TRADEMARK OFFICE**

Applicant(s):	Stavrianopoulos et al.	)	
Serial No.:	08/486,070	)	Group Art Unit: 1631
Filed:	June 7, 1995	)	Ex'r: Ardin H. Marschel, Ph.D.
For:	<b>ARRAYS AND SYSTEMS COMPRISING ARRAYS FOR GENETIC ANALYSES AND OTHER APPLICATIONS</b>	)	

60 Executive Boulevard  
Farmingdale, NY 11735-4716

Mail Stop Non-Fee Amendment  
Commissioner of Patents  
P.O. Box 1450  
Alexandria, VA 22313-1450

**DECLARATION OF DR. DOLLIE M. W. KIRTIKAR UNDER 37 C.F.R. §1.132**

I, Dollie M. W. Kirtikar, hereby declare as follows:

**BACKGROUND**

1. I am presently Senior Scientist for Enzo Biochem, Inc., 60 Executive Boulevard, Farmingdale, New York 11735-4716, having held that position since 1997. Prior to my present position, I was Research Scientist for Enzo, having been first hired in February 1979. During the early 1990s, I held other positions within Enzo or its subsidiaries, including Production Manager and Supervisor of QA/QC. My professional experience is listed on my curriculum vitae attached as Exhibit 1.

2. In terms of my education, I received my Bachelor of Science (B.Sc.) with honors from the University of Bombay, Bombay, India in 1952, graduating with a major in microbiology and a minor in chemistry. In 1957, I received a Master of Science (M.Sc.) from the Seth C.S. Medical College, University of Bombay. In 1967, I was awarded a Doctor of Philosophy (Ph.D) from the University of Kansas. My doctoral thesis was titled "Phenotypic Transformation."

**Enz-7(P)(C3)**

3. From 1969 to 1971, I received a visiting fellowship from the Nucleic Acid Research Foundation of Netherlands, under which I carried out research on enzymes and factors involved in DNA transcription in the Biochemistry Department, University of Netherlands, Croningen, Netherlands. From 1971 to 1977, I was a research associate in the Biochemistry Department, Case Western Reserve University School of Medicine, Cleveland, Ohio. At Case Western, I conducted research on enzymes involved in DNA repair following treatment with cancer-causing physical and chemical agents. From 1977 to 1978, I was a research associate in the Radiology Department, Stanford University Medical Center, Stanford, California, where I conducted research on DNA-repair deficient bacteria. My education and research experience are listed on my CV (Exhibit 1).

4. During my education and research spanning the years 1961 to 1978, I held a number of teaching positions which are listed on my CV (Exhibit 1).

5. I am the author of several scientific publications and the inventor named on several U.S. and foreign patents and patent applications. My scientific publications are listed on my CV (Exhibit 1). Representative issued U.S. patents are listed on my CV (Exhibit 1). I am also a co-inventor on the above-identified U.S. Patent Application Serial No. 08/486,070, filed on June 7, 1995 (hereinafter "the '070 application"). I am familiar with the specification for the '070 application that was filed on June 7, 1995. I have also read the Declaration of Dr. Jannis G. Stavrianopoulos, who is one of my co-inventors on the '070 application.<sup>1</sup> I have additionally read the Office Action mailed on July 2, 2003.

6. As a co-inventor, I am making this Declaration on behalf of and at the request of the assignee.

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<sup>1</sup> I understand that Dr. Stavrianopoulos's Declaration was submitted to the United States Patent and Trademark Office in Applicants' June 17, 2002 Supplemental Amendment T [Their] April 10, 2002 Amendment Under 37 C.F.R. §1.116.

**EXPERIMENTS UNDERLYING THE '070 APPLICATION**

7. In connection with the present invention, my co-inventors and I discovered that nucleic acids could be fixed to non-porous solid supports and remain available for hybridization and detection. In our research and experiments that led to the present invention and the filing of the first application in January 1983, my co-inventors and I investigated different support materials, shapes, surfaces and treatments with respect to fixing nucleic acids to supports in hybridizable form. In at least two separate instances, we constructed an array of different nucleic acids fixed or immobilized to two different kinds of solid supports. Among the different solid supports I investigated prior to 1983 are those listed below in chronological order (most recent at the bottom). Copies of pages from my laboratory notebook are also included herewith as Exhibits 2-9.

<b><u>SUPPORT/SURFACE</u></b>	<b><u>TITLE OF LABORATORY NOTEBOOK PAGE(S)</u></b>	<b><u>EXHIBIT NO.</u></b>
[flat] microscope slides with slots (preprinted slides) <sup>2</sup>	Detection of glucosylated DNA with fluorescent Con-A on slides (microscope slides with slots)	2
glass tubes	Con-A binding to glucosylated DNA treated glass tubes -- Binding of DNA to glass	3
glass tubes	Concentration curve for DNA binding to activated glass tubes (Con A binding to DNA on the glass surface)	4
plastic wells	DNA binding to activated surfaces -- plastic wells treated with epoxy-glue	5
plastic plates	Activated plastic plates -- Detection of glycosylated DNA by con A -- alkaline phosphatase	6
glass tubes	DNA binding to activated glass surface	7
glass fiber filters	Preparation of glass fibre filter	8
glass tubes/slides	Preparation of Silanized glass	9

<sup>2</sup> My laboratory notebook refers to "microscope slides with slots." In common parlance, "slots" may suggest rectangular channels, troughs or indentations. In the context of my experiments, however, "slots" refers to the flat circles that are preprinted onto flat microscope slides. It is known in the art that 'slots' is a synonym for the circles on preprinted slides, also known as cytology slides.

8. Since the focus of our investigations was attachment of nucleic acids to ostensibly inert materials, such as glass and plastic, I focused upon surface treatments for glass and plastic that would permit such binding. Because the shape of these materials is irrelevant to their surface chemistry, I used a variety of differently shaped supports to carry out these experiments, including microscope slides, glass fibers, test tubes, microtitre plates and wells.

**FLAT MICROSCOPE SLIDES WERE USED IN OUR EXPERIMENTS**

9. As indicated above and as shown in my laboratory notebook pages (Exhibits 2 and 9), on at least two separate occasions I fixed nucleic acids to flat microscope slides.

**ARRAYS OF DIFFERENT NUCLEIC ACIDS WERE CONSTRUCTED ON SEPARATE OCCASIONS WITH DIFFERENT SOLID SUPPORTS**

10. As shown in my laboratory notebook pages (Exhibits 2 and 8), I constructed two nucleic acid arrays<sup>3</sup>, one using preprinted microscope slides and the other using glass fiber filters. See the second page of both Exhibits 2 and 8.

A. In the first array (Exhibit 2, second page), T4 and  $\lambda$  DNA were spotted on twelve locations on the same flat microscope glass slide. The flat microscope slide was a preprinted glass slide, also called a cytology slide.

B. In the second array (Exhibit 8, second page), T4 and  $\lambda$  DNA were each spotted three times on the same glass fiber filter. In fact, two such arrays in the form of glass fiber filters were taped to my original laboratory notebook page

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<sup>3</sup> This is true under a restrictive definition of "nucleic acid array" as meaning a plurality of various nucleic acids arranged on a solid support. Under the plain definition – a plurality of (various or identical) nucleic acids arranged on a solid support – more of my experiments could be characterized as involving construction of nucleic acid arrays.

(Exhibit 8, second page) at the time the experiment was performed and both arrays are still taped there.

**MY EXPERIMENTAL WORK IS IN THE '070 APPLICATION AS FILED**

11. The '070 application as originally filed is completely consistent with the experiments discussed above in that it reasonably conveys that (1) the invention resided in the attachment of nucleic acids to the surface of a non-porous solid support; and that (2) the shape of that surface of the non-porous solid support was irrelevant to the invention; and, therefore, (3) that a person skilled in the art at the time the application was filed would not have viewed arrays as being limited to those including wells or depressions because of the irrelevance of the shape of the surface or solid support to which the nucleic acid was bound.

The first conclusion is manifest from page 10 of the '070 application, stating that:

In accordance with the practice of this invention, analytes in a biological sample are . . . directly fixed to a suitable solid support. . . . It is preferred that the solid support to which the analyte is fixed be non-porous.

The second conclusion, that the shape of the solid support is irrelevant, is likewise made manifest on page 10, which states that:

[I]t is preferred that the solid support to which the analyte is fixed be . . . glass, or alternatively, plastic, polystyrene, polyethylene, dextran, polypropylene, and the like.

It will be appreciated that the above cited passage *makes no reference to the shape of the solid support*. Certainly, had the shape been critical, it would have been specified. Indeed, only later in the specification at page 13, as a *preferred embodiment*, solid supports with wells or depressions are cited:

Yet **another** aspect of the method of the present invention involves generating the soluble signal . . . in a device capable of transmitting light therethrough for the detection of the signal by spectrophotometric techniques. . . . **Examples** of the devices useful in the spectrophotometric analysis of the signal included conventional apparatus employed in diagnostic laboratories, i.e., plastic or glass wells, tubes, cuvettes or arrangements of wells, tubes or cuvettes.

Accordingly, point 3 is clearly conveyed, that a person skilled in the art was told by the '070 application that the invention resided in attaching nucleic acids to a solid support without any reference to the shape of that solid support; and that solid supports with wells or depressions were but a preferred embodiment, THEREFORE, the '070 application conveyed that an array as disclosed need not be limited to one with wells or depressions.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that any such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Oct. 28th 2003  
Date

Dollie M. W. Kirtikar  
Dollie M. W. Kirtikar, Ph.D.

\*\*\*\*\*

**Dollie M.W.Kirtikar**  
71-30 162<sup>nd</sup> Street, Unit #3  
Fresh Meadow, NY 11373  
(415) 327-3442

**EDUCATION:**

Bachelor of Science B.Sc.

April 1952 with Honors.  
Microbiology as Major; Chemistry as minor.  
St.Xavier's College, Bombay,India.  
University of Bombay.

Master of Science M.Sc.

February 1957 (three years with research  
Microbiology as major; Biochemistry as minor.  
Seth G.S.Medical College, Eombay, India.  
University of Bombay.  
Thesis: Typing of Salmonella Paratyphi A Strains For All  
Over The World

Doctor of Philosophy Ph.D.

June 1967.  
University of Kansas. Kansas (U.S.)  
Title of Doctorial Thesis: Phenotypic Transformation

**RESEARCH EXPERIENCE:**

March 1969 to March 1971

Visiting fellowship from Nucleic Acid Research  
Foundation of Netherlands  
Biochemistry Department, University of Netherlands at  
Groningen, Netherlands

Molecular biology with emphasis on enzyme, and  
factors involved in DNA transcription

May 1971 to September 1977

Research Associate at Biochemistry Department,  
Case Western Reserve University  
School of Medicine, Cleveland, OH 44106

Molecular biology with emphasis on enzyme  
involved in DNA repair after treatment with physical and  
chemical agents known to cause cancer.

October 1977 to June 1978

Research Associate at Radiology Department  
Stanford University Medical Center,  
Stanford, CA 94305

Molecular biology and genetics with emphasis on DNA-  
repair deficient bacteria

February 1979 to February 1997

Research scientist at Enzo Biochem, Inc. and its  
subsidiary, Enzo Life Sciences, Inc.  
(formerly Enzo Diagnostics, Inc.)



Also held position as Production Manager and  
Supervisor of QA/QC

February 1997 to Present

Senior Scientist at Enzo Life Sciences, Inc.  
(formerly Enzo Diagnostics, Inc.)

**TEACHING EXPERIENCE:**

1961-1963      Teaching Assistantship at Microbiology Department, University School of Kansas  
Medical Center, Kansas City, Kansas.

General laboratory assistance in various courses offered to medical and graduate  
students. Preparation of course material, supervision of students conducting  
experiments, assistance in lab experiments, conferences with students, etc.

1965            Laboratory Instructor in Department of Pathology and Bacteriology at Topiwalla  
National Medical College, Bomoay, India.

Supervision of clinical laboratory staff.  
Training of research personnel such as technicians, graduate students. Conferences  
with students etc. on particular research projects. .

1969-1971      Biochemistry Department, University of Netherlands at Groningen, (March) Netherlands.

Training of graduate students and technicians. Conferences and discussions with  
graduate students on particular research project

1972-1977      Biochemistry Department, Case Western Reserve University School of Medicine,  
Cleveland, OH 44106

General supervision of laboratory personnel; training and supervision of technicians,  
graduate and college students; initiation of new personnel in the laboratory group in  
various techniques. Conferences and discussions and suggestions to various personnel  
about specialized techniques in molecular biology.

Jan. 1978 to      Radiology Department, Stanford University Medical Center, Stanford, CA 94305  
June 1978

Training and supervision of one technician in molecular genetics with particular  
emphasis on mapping of bacterial mutants. Preparation of phage lysate, transduction  
and conjugation expts, construction of strains etc.

#### PUBLICATIONS:

1. Mridula W. Kirtikar, Typing of Salmonella Paratyphi A Strains For All Over The World. *J. Post Graduate Medicine of India* (1957)
2. Mridula W. Kirtika, D.D. Banker, and N.W. Purandare, Bacteriophage Typing of 371 Strains of Salmonella Paratyphi A. *J. Post-Grad. Med., Vol. VI, 6* (1960)
3. N.Veeraraghavan and M.W. Kirtikar, Biological Characteristics of Influenza virus Strains Isolated at the Government of India Influenza Centre, Coonoor, during 1950-69, *Bull. World Hlth.Org., 24, 687* (1961)
4. N.Veeraraghavan, M.W. Kirtikar and T. Sreevalsan, Studies on the Cultivation of Influenza Virus in vitro. *Bull. World Hlth. Org., 24, 711* (1961)
5. Mridula W. Kirtikar, Phenotypic Transformation. *Dissertation Abs Intl., 28 (7), 2708* (1967)
6. Mridula W. Kirtikar and Jacob D. Duerksen, A Penicillinase-Specific Ribonucleic Acid Component from Bacillus Cereus. I. Ribonucleic Acid Extraction and Definition of the in Vivo Test System. *Biochemistry, 7, 1172* (1968)
7. Mridula W. Kirtikar and Jacob D. Duerksen, A Pencillinase-Specific Ribonucleic Acid Component from Bacillus Cereus. II. Partial Characterization of the Active Component. *Biochemistry, 7, 1183* (1968)
8. Dollie M. W. Kirtikar and Akira Kaji, Stimulation of Phage Ribonucleic Acid-dependent Incorporation of Amino Acids by 5 S Ribonucleic Acid. *J.Biol.Chem., 243, 5345* (1968)
9. Shelk-Mumtaz Hadi, D. M. Kirtikar, and D.A. Goldthwait, (Endonuclease II of Escherichia coli. Degradation of Double- and Single-Stranded Deoxyribonucleic Acid. *Biochemistry, 12, 2747* (1973)
10. D.M. Kirtikar and D.A. Goldthwait, The Enzymatic Release of O<sup>6</sup>-methylguanine and 3-methyladenine from DNA Reacted with the Carcinogen N-methyl-N-nitrosourea. *Proc. Nat. Acad. Sci., 71, 2022* (1974)
11. D.A. Goldthwait, D.M. Kirtikar, S.M.Hadi, and E.C. Friedberg, Molecular Mechanisms for repair of DNA, Part A. *Plenum Publishing Company, p.191-196* (1975)
12. D.M. Kirtikar, J. Slaughter and D.A. Goldthwait, Endonuclease II of Escherichia coli: Degradation of  $\gamma$ -Irradiated DNA. *Biochemistry, 14, 1235* (1975)
13. Dollie M. Kirtikar, Anthony Dipple and David A. Goldthwait, Endonuclease II of Escherichia coli: DNA Reacted with 7-Bromomethyl-12-methylbenz[a]anthracene as a Substrate. *Biochemistry, 14, p.5548* (1975)
14. D.M. Kirtikar, J.P. Kuebler, A. Dipple, and D.A. Goldthwait, Endonuclease II of Escherichia coli and Related Enzymes, Fundamentals in Cancer Prevention: 6th Int. Symp. of the Princess Takamatsu Cancer Research Fund. *University of Tokyo Press, p.349* (1975)

15. D.M. Kirtikar, J.P.Kuebler, A. Dipple, and D.A. Goldthwait, Enzymes Involved in Repair of DNA Damaged By Chemical Carcinogens and  $\gamma$ -Irradiation. *Cancer Enzymology* (eds. J. Schultz and F.E. Ahmed, *Eighth Miami Winter Symposium, Academic Press, p.139* (1976)
16. Dollie M. Kirtikar and David A. Goldthwait, Endonuclease II and the Apurinic Acid Endonuclease of E. Coli. *Fed. Proc.*, 35, 1589 (1976)

**PATENTS:**

U.S.

- o 5,241,060
- o 5,260,433
- o 4,994,373

\* \* \* \* \*

Detection of glycosylated DNA  
with fluorescent con-A.  
on slides.

6/1/82.

Norman  
Keller  
6/1/82

Microscope slides with slots

Treated for 5' with 0.1% HCL

Washed with water 5X

Treated with subbing solution

0.1% CrK(SO<sub>4</sub>)<sub>2</sub>

1% gelatin

for 5 minutes

Air dried.

Apply DNA 20  $\lambda$

Air Dry

Wash with 70% EtOH 20'

followed by 95% EtOH 20'

Air Dry

Apply con A 20  $\lambda$  in NaCl-Mg<sup>++</sup> 10 mM each

~~Air~~ + Keep in <sup>HUMID</sup> Hybrid chamber  
for 30'

Rinse with PBS

Keep in PBS 7 minutes

20λ 5λ 10λ 15λ 20λ DNA

DNA (T<sub>4</sub> or λ)

20λ 5λ 10λ 15λ 20λ

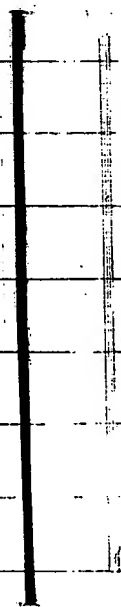
con A - 20λ 20λ 20λ 20λ 20λ

all positive\* 0 0 0 0 0 0 T<sub>4</sub> DNA

all negative 0 0 0 0 0 0 λ DNA

Positive

characteristic  
\* gave a fluorescent green color



Wash slides 0.1N HCl, 5min  
 Rinse d H<sub>2</sub>O 5X  
 10% gel, 0.1% K<sub>2</sub>Cr<sub>2</sub>O<sub>7</sub> 5min/dry

0.5

Apply Chromosome

70% EtOH, 95%, dry

0.1N NaOH

EtOH

95% Form 68°C, 1hr

EtOH

Hybridize

45% form, 4x22, 60-65°C

CONA 50x/wt 20x

20x

T4 DNA

○ ○ ○ ○ ○ ○

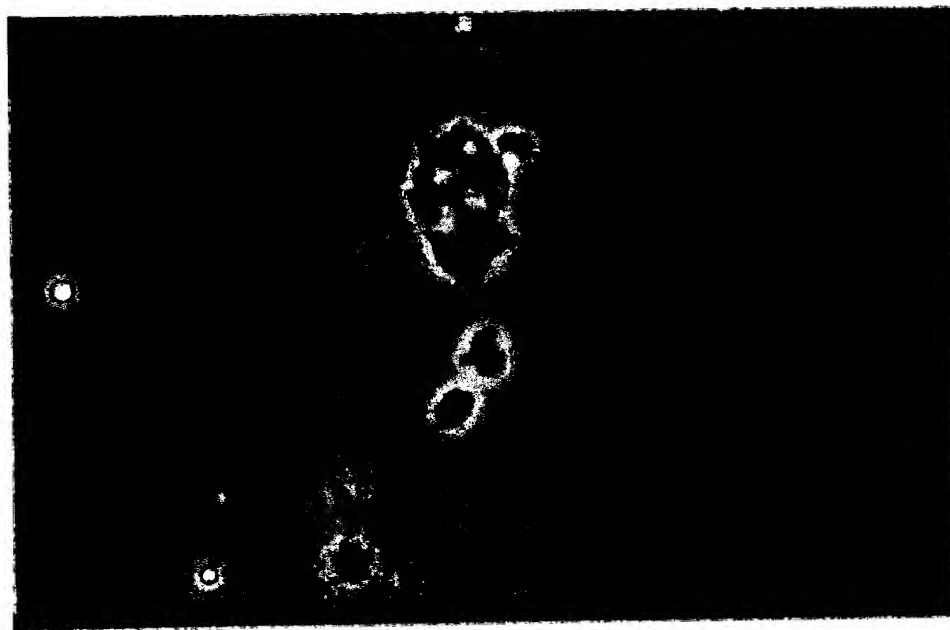
λ DNA

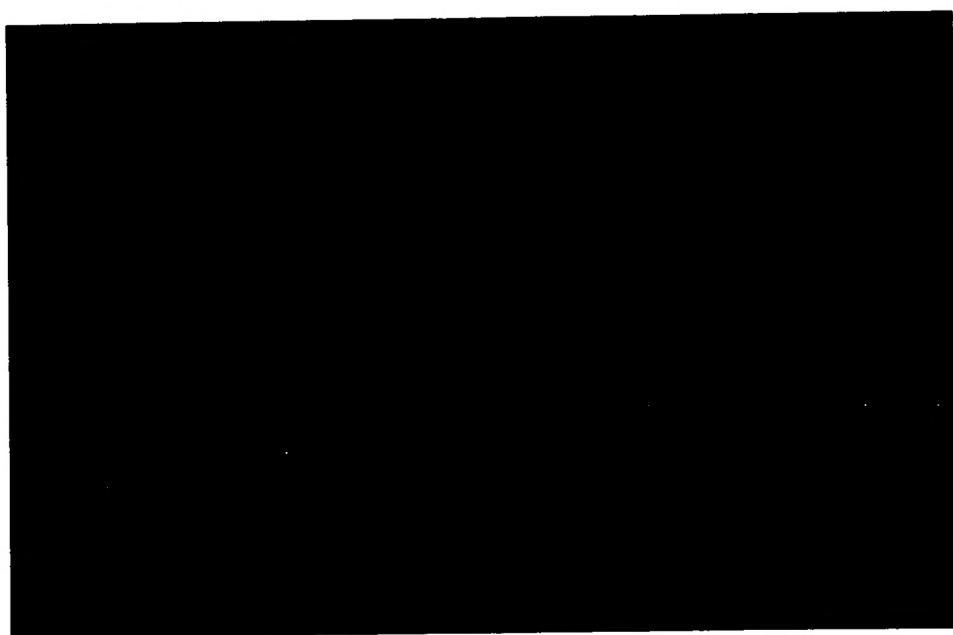
○ ○ ○ ○ ○ ○

CONA

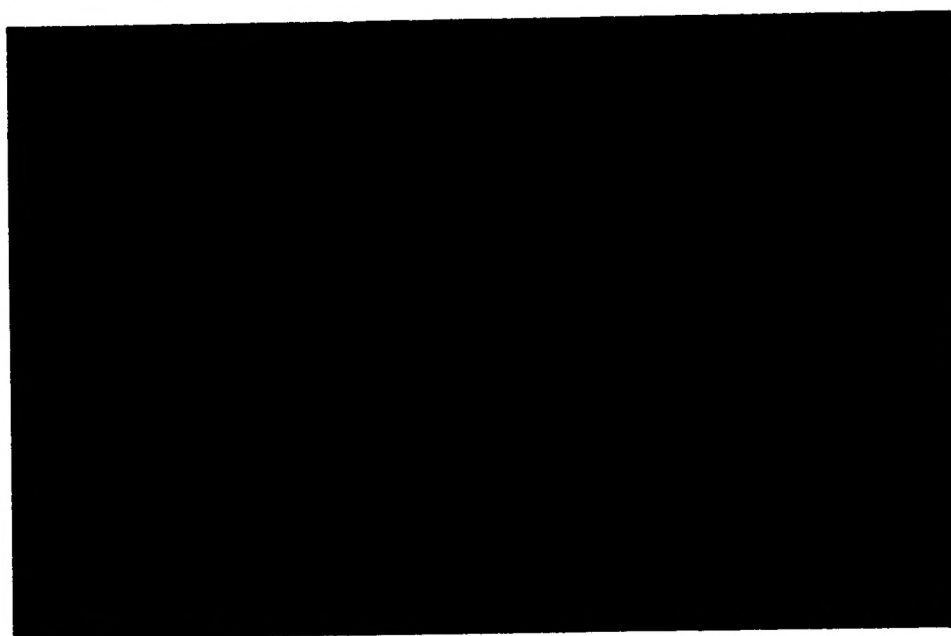
-

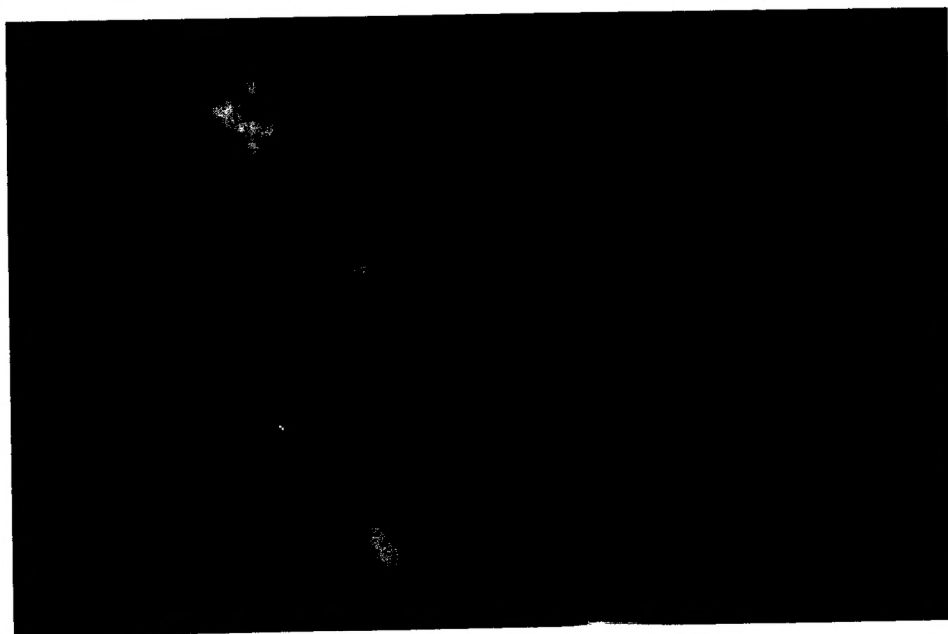
20











6.13.82

# Con A binding to glucosylated DNA

Treated glass Tubes

Norman Kelly

6/13/82

	1	2	3	4	5	6	7	8	9
T <sub>4</sub> DNA	100 $\lambda$	-	100 $\lambda$	-	100 $\lambda$	-	-	-	-
CT DNA	-	100 $\lambda$	-	100 $\lambda$	-	100 $\lambda$	-	-	-
Fluorescein A	-	-	100 $\lambda$	100 $\lambda$	100 $\lambda$	100 $\lambda$	100 $\lambda$	100 $\lambda$	-
carrot phosphatase	-	-	-	-	+	+	+	-	+

After each addition tubes incubated at RT for 15-30 minutes

Washed repeatedly  $\bar{c}$  PBS 100  $\lambda$  aliquots 20 times.

Tubes 1 and 2 stained with 100  $\lambda$  of 1  $\mu$ g/ml EtBr in PBS 30' in the dark. at room temp

To tubes 3, 4 and 8 added con A

fluorescent 5  $\mu$ g/ml in NaCl Mg<sup>++</sup> made up to 10 mM

Left at room temp for 1 hour

washed repeatedly  $\bar{c}$  PBS.

## Binding of DNA to glass

checked under UV lamp after Ethidium bromide staining of tubes 1 + 2 and Fluorescent cou A interaction with tubes 3, 4 + 8



Tubes 1 + 2 gave a red fluorescence characteristic of Ethidium bromide and tube 3 gave a green fluorescence. ~~the~~  
Tubes 4 and 8 gave a ~~red~~ visible green fluorescence.

To tubes 5, 6, 7 added unlabelled cou A  
5  $\mu\text{g}/\mu\text{l}$  in PB containing NaCl,  $\text{Mg}^{2+}$ ,  $\text{Ca}^{2+}$   
left at room Temp for 60 minutes  
washed repeatedly with PBS 20 times.  
washed with Imidazole-HCl 5x 0.2M pH 6.5  
Added acid phosphatase 1/10 diluted 10x 2 units  
Added diluent containing BSA 100  $\lambda$   
left at RT for 30'  
Washed repeatedly with Imidazole-HCl 20 times.  
Added 20  $\lambda$  substrate  
80  $\lambda$  buffer

left at RT

color intense yellow in tube #5

immediately Added 1.0 ml bicarbonate  
checked at 4 min.

$A_{300}$  in 50' on 1:1 diluted samples.

# Tube  $A_{300}$  vs  $H_2O$  blank.

5	0.498
6	0.065
7	0.055
9	0.011

$\lambda$  DNA nick-translated  $\bar{c}$  malto-triose substituted  
UTP was attached to the glass similarly  
Con A was added, washed off.

detection of conA binding to DNA  
was carried out using  
acid phosphatase in the same  
manner as for T4 DNA.

Tube #		$A_{300}$
10	$\lambda$ DNA control nick translated $\bar{c}$ UTP	0.005
11	$\lambda$ DNA -test (nick translated with $\bar{c}$ malto-triose UTP)	0.368
12	con A + acid phosphatase	0.012
13	acid phosphatase alone.	0.008

Norman K. 6/17/82  
6.17.82

# Concentration curve for DNA binding to activated glass tubes.

$T_4$  DNA -  $^3H$   $T_4$  DNA nick-translated using  
3.29 micrograms/ml  $^3H$  dATP  
 $\lambda$  DNA -  $^3H$   $\lambda$  DNA nick-translated using  
3.615 micrograms/ml  $^3H$  dATP

$5\lambda$   $^3H$   $T_4$  DNA 16.45 ng  $^3H$ cpm  
7809 } 7220  
6630  
approx. 439 cpm/nanogram

$5\lambda$   $^3H$   $\lambda$  DNA 18.075 ng  $^3H$ cpm  
5044 } 4699  
4354  
approx. 260 cpm/nanogram

$^3H$  6.17.82  
 $T_4$   $5\lambda$  16.45 ng  $^3H$   $\lambda$  DNA 6.17.82  
 $5\lambda$  18.075 ng

1 01	001.00	1 05	001.00
	003.00		003.00
	007.00		007.00
1 02	001.00	1 06	001.00
	003.00		003.00
	006.00		006.00

4699

$^3\text{HT}_4$  DNA 10 $\lambda$  diluted to 1.0 ml.

32.9 ng/ml.

DNA

~~10~~

- |      |                                      |                      |
|------|--------------------------------------|----------------------|
| 1. A | 10 $\lambda$ + 490 $\lambda$ buffer  | - 658 picograms/ml   |
| 2. B | 20 $\lambda$ + 480 $\lambda$ buffer  | - 1.316 nanograms/ml |
| 3. C | 40 $\lambda$ + 440 $\lambda$ buffer  | - 2.632 nanograms/ml |
| 4. D | 50 $\lambda$ + 450 $\lambda$ buffer  | - 3.290 nanograms/ml |
| 5. E | 100 $\lambda$ + 400 $\lambda$ buffer | - 6.58 nanograms/ml  |
| 6. F | 200 $\lambda$ + 300 $\lambda$ buffer | - 13.16 nanograms/ml |
| 7. G | 500 $\lambda$ + 0                    | - 32.9 nanograms/ml. |

$^3\text{HT}_4$  DNA 25 $\lambda$  diluted to 1.0 ml

82.25 ng/ml.

- |      |                          |              |
|------|--------------------------|--------------|
| 8. H | 500 $\lambda$ + 0 buffer | 82.25 ng/ml. |
|------|--------------------------|--------------|

$^3\text{HT}_4$  DNA 50 $\lambda$  diluted to 1.0 ml.

164.5 ng/ml.

- |      |                          |              |
|------|--------------------------|--------------|
| 9. I | 500 $\lambda$ + 0 buffer | 164.5 ng/ml. |
|------|--------------------------|--------------|

$^3\text{HT}_4$  DNA 100 $\lambda$  diluted to 1.0 ml.

- |       |                           |              |
|-------|---------------------------|--------------|
| 10. J | 500 $\lambda$ + 0 buffer. | 329.0 ng/ml. |
|-------|---------------------------|--------------|

100  $\lambda$  of each solution checked for radioactivity  
after dilution  $\bar{c}$  200  $\lambda$  of 5 mM Tris 7.4

100  $\lambda$  of each solution placed in  
A nonactivated glass Tubes  
B activated glass Tubes.

After 5' at R.T. solutions removed  
carefully

tubes washed 2 x  $\bar{c}$  100  $\lambda$   
portions of PBS-Mg<sup>++</sup>

~~All~~ ~~con~~ washes and originals combined  
and counted.

<sup>3</sup>H epm

	Original - Bkg(3.5)			Non-Activated - Bkg(3.5)		Activated - Bkg(3.5)	
①	65.8 pgm	23	100%	23	100%	16	69.56%
②	131.6 pgm	32	100%	24	77.5%	27	84.38
③	263.2 pgm	48	100%	49	100%	32	66
④	329.0 pgm	65	100%	70	107.6%	39	60
⑤	658 pgm	106	100%	111	104.7%	97	91.5
⑥	1.316 ngm	245	100%	255	104%	167	68.16
⑦	3.29 ngm	527	100%	569	108%	468	82.25
⑧	8.225 ngm	1337	100%	1348	100.8%	979	73
⑨	16.45 ngm	2387	100%	2428	101.7%	2089	87.5%
⑩	32.9 ngm	4053	100%	3883	95.8%	3060	75.5



oohi 6/21/82

2'

Remained on Activated  
glass surface

1. 06  
102.00  
100.00  
10007.00  
1. 07  
107.00  
100010.00  
101.00  
1. 04

Volle  
6/21/82

105

34  
5878

70

Q

9

Q

1

1. 20 picograms.

2. 20,56

3. 89.49

4. 131.6

5, 55, 93

6. 419.00

7. 584.00

8. 2220.75

9. 2052.96

10 8060.5

10'

picograms on glass surface

1 28.95

2. 17.10

3. 131.6

4. 118.44

S. 184

6. 355

7. 428

8 2303

9. 2139

10. 855-4

# Con A binding to DNA on the glass surface

Con A 100  $\mu\text{g}/\text{ml}$  in PBS

1  $\mu\text{g}/\text{ml}$  "

10  $\text{ng}/\text{ml}$  "

DNA on Surface : Con A R.T. 60'

1	<del>20 pg</del> 28.95	100 pg
2	<del>20.56</del> 17.10	100 pg
3	<del>89.49</del> 131.6	100 ng
4	<del>131.6</del> 118.44	100 ng
5	<del>55.93</del> 184	100 ng
6	<del>419</del> 355	10 $\mu\text{g}$
7	<del>584</del> 428	10 "
8	<del>2220</del> 2303	10 "
9	<del>2052</del> 2139	10 "
10	<del>8060</del> 8554	10 "

6.30.82.

Con A removed and the tubes washed

5 times with PBS. 100  $\lambda$

2 times with Imidazole Buffer pH 6.5

Alkaline phosphatase 10  $\text{ng}/\text{ml}$  diluted

to 0.5  $\text{ng}/\text{ml}$  in Tris pH 7.4

100  $\lambda$  added to each tube.

Incubated at RT 30 minutes.

Removed and washed with Imidazole Buffer <sup>pH 6</sup>

3 times 100  $\lambda$

Added 20 microliters of substrate 0.1 M

80 microliters of Tris pH 7.4 ~~500~~ 2 M

Incubated at RT.

Within 10 min. the color develops.

After 30 min. at RT.

Diluted 1/10 in 1 M Tris pH 7.4

and read at 410 vs. substrate blank.

$A_{410}$

$\times 10$   ~~$\times 10$~~

DNA on surface.

1	28.95 pcg.	0.385
2	17.10	0.296
3.	131.6	0.6
4.	118.44	0.58
5.	184	0.6
6.	355	0.67
7.	428	0.43
8.	2303	<del>0.68</del>
9.	2139	<del>0.68</del>
10.	8559	<del>0.69</del>

untreated 11. Enzyme + Substrate Blank. 0.71

untreated 12. Enzyme + Substrate Blank + con A 0.705

treated 13. Enzyme + Substrate + con A Blank. 0.06

treated 14. Enzyme + Substrate Blank. 0.03

6.30.82

DNA binding to activated surfaces.

plastic wells treated with epoxy-glu.  
5 $\lambda$  and 10 $\lambda$   
air-dried.

<sup>3</sup>H T<sub>4</sub> nick translated DNA 32.9  $\mu$ g/ml  
diluted to contain

A	0.658	$\mu$ g/ml.	in	5mM	TRIS	7.4.
B	1.316	"				
C	2.632	"				
D	3.290	"				
E	6.580	"				
F	13.160	"				
G	32.90	"				

50 microliter added per well.

after 10' at RT.

samples removed

wells washed with PBS pH 6.5 twice.

50 microliter aliquots.

all washes combined and counted

Samples

50  $\lambda$  of 1  $\mu$ g/ml con A in PBS. 19<sup>th</sup>  
applied to each well.

Left at RT. for 30'

washed  $\bar{a}$  in diazole buffer twice

Applied 50 microliters of alkaline phosphatase  
0.5 mg/ml

After 30 minutes at RT.

washed in Tris containing 0.15 M KCl  
3 times.

finally with Tris alone.

Applied 0.1 M substrate: 5  $\lambda$  in  
45  $\lambda$  Tris 7.4 2.0 M.

Immediate color developed. but  
the enzyme by itself also sticks.

6.20.82

	1	2	3	4	5	6	7	8	9	
1	A	Epoxy Glu	5x							
2										
3										
4										
5	DNA	32.9	65.8	131.6	164.5	329	658	1645		
6	Applied.	← pico grams →								
7		14	29.6	59.2	74	148	296	740.3		
8		← 3H epm →								
9										
10										
11	DNA									
12	not	13.5	13.5	24	49	73	165.5	393.5		
13	bound.	← 3H epm →								
14										
15										
16	DNA									
17	on	0.5	16.1	35.2	25	75	130.5	346.8		
18	surface	← 3H epm →								
19										
20										
21		1.11	35.74	78.15	55.5	166.5	289.7	769.9		
22		← pico grams →								
23										
24										
25	Con A	50	50	50	50	50	50	50	50	
26		← nanograms →								
27										
28										
29	Alk. Phosp.	25	25	25	25	25	25	25	25	25
30		← micrograms →								
31										
32										
33										
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6.30.82.

[illegible]

Dollie  
6/29/82

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000434.00  
1 34

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000434.00



Norman K. ...  
7/13/82

Activated plastic plates

7.13.82

Detection of glycosylated DNA

Norman K. ...

by con A - alkaline phosphatase

7/13/82

$^3\text{H T}_4$  DNA

wick translated 3.29  $\mu\text{g}/\text{ml}$

diluted in PBS buffer to

32.9 nanograms/ $\mu\text{ml}$ .

Con A

10  $\mu\text{g}/\text{ml}$  in 2.0 M NaCl

50  $\mu\text{g}$  in 5  $\lambda$  lambda.

Alkaline phosphatase

0.5  $\mu\text{g}/\text{ml}$  in

5 mM TRIS pH 7.4

4-Nitrophenyl phosphatase

1  $\mu\text{g}/\text{ml}$  in

1 M TRIS pH 7.4

BSA

20  $\mu\text{g}/\text{ml}$  phosphatase from Janos.

1. DNA 50  $\lambda$  RT 10'

Rinse 3 x  $\bar{e}$  100  $\lambda$  PBS

Count

2. Con A 5  $\lambda$  (50  $\mu\text{g}$ ) in PBS  $\text{Mg}^{++}$

Total 50  $\lambda$

RT 30'

3. Rinse 3 x  $\bar{e}$  PBS  $\text{Mg}^{++}$  100  $\lambda$  each rinse

2 x  $\bar{e}$  TRIS.HCL 100  $\lambda$  each rinse

4. BSA 20  $\mu\text{g}/\text{ml}$  50  $\lambda$

37° 2 hours.

5. Rinse  $\bar{e}$  TRIS.HCL 5 mM 2 x

6. Alkaline phosphatase 50  $\lambda$  55  $\mu\text{g}$

30' RT.

Winnam Keller  
7/13/82

7. Riuse 5 x e 100  $\lambda$  of 0.3 M NaCl

8. Substrate 2 mg/ml 80x 20  $\mu$ g

Incubate at 37°C

# MICROELISA

	A	B	C	D	E	F	G	H
DNA	+	+	+	+	-	-	-	-
ConA	+	+	-	-	+	+	-	-
BSA	+	+	+	+	+	+	+	+
AP	+	-	+	-	+	-	+	-
NaCl	+	+	+	+	+	+	+	+
Substrate	+	+	+	+	+	+	+	+

1 A	>	1.007
1 B		0.002
1 C	>	0.749
1 D		0.006
1 E	>	0.727
1 F		0.025
1 G		0.149
1 H		0.000
2 A	>	1.379
2 B		0.001
2 C		OVER
2 D		0.064
2 E		OVER
2 F		0.026
2 G		0.015
2 H		0.012

7/13/82  
1.00  
0.05.00  
0.01.00 Bkg.  
0.0017.20  
1.00  
0.05.00 origini  
0.01.00  
0.0040.00  
1.00  
0.05.00 well  
0.01.00 ①  
0.0013.00  
1.01  
0.05.00 well  
0.01.00 ②  
1.00  
0.05.00 well  
0.01.00 ③  
0.0022.00  
1.00  
0.05.00 well  
0.01.00 ④  
0.0013.00

A<sub>410</sub>

A	0.149
B	0.000
C	0.015
D	0.012
E	0.099
F	0.008
G	0.053
H	0.000

3 A		OVER
3 B	>	0.304
3 C		OVER
3 D	>	0.343
3 E		OVER
3 F	>	0.401
3 G		0.099
3 H		-0.001
4 A		OVER
4 B		0.002
4 C	>	0.381
4 D		0.012
4 E	>	0.830
4 F		0.031
4 G		0.000

8.2.82.

[illegible]

[illegible]

	1	2	3	4	5	6	7	8	9	
1		Add	20 $\mu$	of	0.1 M	substrate				
2			80 $\mu$	of	0.2 M	Tris-glycine buffer (pH 8.5)				
3										
4										
5										
6										
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Incubate at 37° for one hour.

~~Dilute~~

Stop Rx @ 2.0 ml 5% NaHCO<sub>3</sub>.

Check OD at A<sub>410</sub>.

	1	2 DNA	3 Cou A	4 Ficoll	5 Cou A	6 Acid P.	7	8 $\Delta$ 410	9
1									
2									
3	A	50 ng	+	-	-	+		0.285 x 20	
4									
5	A <sub>2</sub>	50 ng	+	+	+	+		0.35 x 20	
6									
7	B <sub>1</sub>	5 ng	+	-	-	+		0.175 x 20	
8									
9	B <sub>2</sub>	5 ng	+	+	+	+		0.23 x 20	
10									
11	C <sub>1</sub>	500 peg	+	-	-	+		0.06 x 20	
12									
13	C <sub>2</sub>	500 peg	+	+	+	+		0.12 x 20	
14									
15	D <sub>1</sub>	50 peg	+	-	-	+		0.11 x 5	
16									
17	D <sub>2</sub>	50 peg	+	+	+	+		0.22 x 5	
18									
19	E <sub>1</sub>	20 peg	+	-	-	+		0.144 x 2	
20									
21	E <sub>2</sub>	20 peg	+	+	+	+		0.208 x 2	
22									
23	F <sub>1</sub>	-	-	-	-	+		0.006	
24									
25	F <sub>2</sub>	-	+	+	+	+		0.007	
26									
27	G <sub>1</sub>	-	-	-	-	+		0.006	
28									
29	G <sub>2</sub>	-	-	+	-	+		0.005	
30									
31	Control	-	-	-	-	+		0.35 x 20	

Preparation of glass fibre filters.

EGF Filters. soaked in 5% Nitric Acid  
For 4 days.

Washed in  $H_2O$

DRIED 24 hours at  $100^\circ$

10% Organosilane prepared pH 3.11  
adjusted in 6N HCL [and end point in  
diluted HCL]

Equilibrated to  $75^\circ$  in brown bottle.

Washed filters dropped one by one in  
solution.

Swirled gently once so all the filters  
were evenly dispersed.

Incubation at  $75^\circ$  for 2 hrs and 45  
minutes.

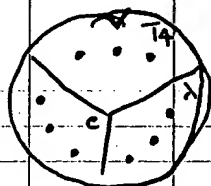
Filters removed liquid sucked off on  
vacuum filtration apparatus.

Filters washed once with distilled water

DRIED o/n at 100°

8/28/82.

Two Filters used



T<sub>4</sub> DNA 1λ per spot 0.2 μg/μl

λ DNA 1λ per spot 0.2 μg/μl

PBS.Mg<sup>++</sup> 1λ per spot

AIR DRY.

Wash  $\bar{c}$  PBS.Mg<sup>++</sup> on Filtration apparatus.

Filter # 1 stained  $\bar{c}$  E+BR.

Filter # 2 soaked in FITC-labelled con A  
0.5 μg/μl 100 λ per filter.

excess fluid removed by filtration

Filter washed with PBS.Mg<sup>++</sup> buffer.





9/1/82

	1	2	3 <sup>3</sup> HT <sub>4</sub>	4	5 CTDNA 2.5 µg/ml	6	7 X <sub>1</sub> SSC	8 3.0 M NaOH	9 2.0 M Am Ac
1									
2									
3									
4		0	20 λ		0		160 λ	20 λ	200 λ
5									
6									
7		1	"		10 λ		150 λ	20 λ	200 λ
8									
9									
10		2	"		20 λ		140 λ	20 λ	200 λ
11									
12									
13		3	"		40 λ		120 λ	20 λ	200 λ
14									
15									
16		4	"		80 λ		80 λ	20 λ	200 λ
17									
18									
19		5	"		160 λ		0	20 λ	200 λ
20									
21									
22									
23		2	Sets.						
24									
25									
26									
27									
28									
29		One	set	filtered thru			activated glass filters		
30									
31		Control	set	filtered thru			EGF untreated		
32									
33									
34									
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Radioactivity checked.

1 23  
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001.50  
018086.00  
1 24  
002.00  
000.00  
000019.00  
1 25  
002.00  
007.00  
000647.50  
1 26  
002.00  
015.00  
000163.50  
1 27  
002.00  
010.00  
000223.50  
1 28  
002.00  
015.00  
000124.50  
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000117.00  
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000155.00  
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000018.00  
1 32  
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000115.00  
1 33  
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002.00  
020.00  
000054.00  
1 35  
002.00  
020.00  
000080.50  
1 36  
002.00  
020.00  
000055.00  
1 37  
002.00  
000.00  
000032.00

Treated  
Filters

EFFICIENCY LINE® 22-205



	1 $^3\text{H}$ cpm	3 Competition	5	Counts on Filter	% Inhibition	Capa of Fil
1	T4 DNA	unlabelled				
2		CT DNA				
3		ug.				
4						
5						
6	18086	0		18035		
7						
8		25		18063		
9						
10		50		18023		
11						
12		100		18024		
13						
14		200		18017		
15						
16		400		18055		
17						
18						
19						
20						
21						
22						
23						
24	Untreated					
25	Filters					
26						
27						
28	18086	0		18155		
29						
30		25		18018		
31						
32		50		17115		
33						
34		100		8053		
35						
36		200		3723		
37						
38		400		2413		
39						
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50						

## Preparation of Silanized glass :

Glass Tubes (slides) are boiled for 45 min. in 5% nitric acid solution, then washed and dried for 24 hours at  $115^{\circ}\text{C}$ . The surface area to be activated is dipped into a 10% solution of gamma-aminopropyl-triethoxysilane (Union Carbide Silicones A-1600) dissolved in distilled water and pH adjusted to 3.45 with ~~the~~ normal hydrochloric acid. The suspension is placed for 2.75 hours at  $75^{\circ}$ , removed and then washed with the ~~same amount of~~ <sup>same amount of</sup> water ~~(same volume)~~ <sup>(same volume)</sup> twice and then dried overnight at  $100^{\circ}\text{C}$ . The resulting material has an available alkylamine.

Activated glass tubes were used to show con lectin-binding to glucosylated [glucose-substituted nick translated] DNA. Experimental data indicated that ~~although glucosylated and non~~ although there was no difference in binding of DNA either glucosylated

⊕ the endpoint of the pH titration of  
made with .6M HCl (1/10 dilution)  
the solution warms up during the  
neutralization so that the pH ~~at~~  
reads 3.2 which corresponds to 3.4V  
at 25°C.

or non-glucosylated DNA to activated glass surface [range 60 picograms to 10 nanograms as detected by  $[^3H]$  radioactive DNA], fluorescent labelled con-A bound only to glucosylated DNA [DNA was counter stained with ethidium bromide]. ~~Further experiments indicated that~~ That lectin bound only to glucosylated [glucose-substituted DNA by nick translation] was further substantiated by the fact that ~~horse~~ <sup>gold</sup> ~~phosphatase~~ ~~radish peroxidase~~  $\rightarrow$  (a glycoprotein) bound ~~only~~ to con A which already was stuck to glucosylated DNA. Both ~~con A~~ Enzyme activity was ~~found~~ not washed off from these tubes. Both con A and horse radish peroxidase do not bind to the activated glass surface.